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				* Walcome to STN International * * * * * * * * *
				* Welcome to STN International * * * * * * * * *
NEWS	1			Web Page for STN Seminar Schedule - N. America
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NEWS	3	MAR	31	CAS REGISTRY enhanced with additional experimental
				spectra
NEWS	4	MAR	31	CA/CAplus and CASREACT patent number format for U.S. applications updated
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NEWS	10	APR	28	IMSRESEARCH reloaded with enhancements
NEWS	11	MAY	30	INPAFAMDB now available on STN for patent family
				searching
NEWS	12	MAY	30	DGENE, PCTGEN, and USGENE enhanced with new homology
			0.5	sequence search option
NEWS		JUN		EPFULL enhanced with 260,000 English abstracts
NEWS NEWS		JUN		KOREAPAT updated with 41,000 documents USPATFULL and USPAT2 updated with 11-character
NEWS	13	JUN	13	patent numbers for U.S. applications
NEWS	16	JUN	19	CAS REGISTRY includes selected substances from
112110		0 011		web-based collections
NEWS	17	JUN	25	CA/CAplus and USPAT databases updated with IPC
				reclassification data
NEWS	18	JUN	30	AEROSPACE enhanced with more than 1 million U.S.
				patent records
NEWS	19	JUN	30	EMBASE, EMBAL, and LEMBASE updated with additional
				options to display authors and affiliated organizations
NEWS	20	JUN	20	STN on the Web enhanced with new STN AnaVist
MEMP	20	0.014	30	Assistant and BLAST plug-in
NEWS	21	JUN	3.0	STN AnaVist enhanced with database content from EPFULL
NEWS		JUL		CA/CAplus patent coverage enhanced
NEWS		JUL		EPFULL enhanced with additional legal status
				information from the epoline Register
NEWS	24	JUL	28	IFICDB, IFIPAT, and IFIUDB reloaded with enhancements
NEWS	25	JUL	28	STN Viewer performance improved
NEWS	26	AUG	01	INPADOCDB and INPAFAMDB coverage enhanced
NEWS	27	AUG	13	CA/CAplus enhanced with printed Chemical Abstracts
				page images from 1967-1998
NEWS		AUG		CAOLD to be discontinued on December 31, 2008
NEWS	29	AUG	15	CAplus currency for Korean patents enhanced
NEWS	EXP	RESS	JUNE	E 27 08 CURRENT WINDOWS VERSION IS V8.3,

AND CURRENT DISCOVER FILE IS DATED 23 JUNE 2008.

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FILE 'REGISTRY' ENTERED AT 14:46:40 ON 18 AUG 2008 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2008 American Chemical Society (ACS)

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38 39
ring nodes :
1 2 3 4 5 6 7 8 9 13 14 15 16 17 18
chain bonds :
1-33 2-19 3-34 4-11 8-28 9-27 11-12 12-13 12-35 12-36 14-26 15-39 16-38
17-37 18-25 19-20 19-21 21-22 22-23 22-31 22-32 23-24 23-29 23-30
ring bonds :
1-2 1-6 2-3 3-4 4-5 5-6 5-7 6-9 7-8 8-9 13-14 13-18 14-15 15-16 16-17
17-18
exact/norm bonds :
1-2 1-6 2-3 3-4 4-5 4-11 5-6 5-7 6-9 7-8 11-12 19-20 19-21 21-22
23-24
exact bonds :
1-33 2-19 3-34 8-9 8-28 9-27 12-13 12-35 12-36 14-26 15-39 16-38 17-37
18-25 22-23 22-31 22-32 23-29 23-30
normalized bonds :
13-14 13-18 14-15 15-16 16-17 17-18
isolated ring systems :
containing 1 : 13 :
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11 12 19 20 21 22 23 24 25 26 27 28 29 30 31 32 33 34 35 36 37

chain nodes :

Match level: 1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 11:CLASS 12:CLASS 13:Atom 14:Atom 15:Atom 16:Atom 17:Atom 18:Atom 19:CLASS 20:CLASS

21:CLASS 22:CLASS 23:CLASS 24:CLASS 25:CLASS 26:CLASS 27:CLASS 28:CLASS 29:CLASS 30:CLASS 31:CLASS 32:CLASS 33:CLASS 34:CLASS 35:CLASS 36:CLASS

37:CLASS 38:CLASS 39:CLASS

STR

L1 STRUCTURE UPLOADED

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* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

Structure attributes must be viewed using STN Express query preparation.

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SAMPLE SEARCH INITIATED 14:46:59 FILE 'REGISTRY'
SAMPLE SCREEN SEARCH COMPLETED - 1 TO ITERATE

100.0% PROCESSED 1 ITERATIONS 0 ANSWERS
SEARCH TIME: 00.00.01

FULL FILE PROJECTIONS: ONLINE **COMPLETE**
BATCH **COMPLETE**

PROJECTED ITERATIONS: 1 TO 80 PROJECTED ANSWERS: 0 TO 0

L2 0 SEA SSS SAM L1

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FULL SEARCH INITIATED 14:47:03 FILE 'REGISTRY'
FULL SCREEN SEARCH COMPLETED - 51 TO ITERATE

100.0% PROCESSED 51 ITERATIONS 3 ANSWERS SEARCH TIME: 00.00.01

L3 3 SEA SSS FUL L1

=> file caplus

 COST IN U.S. DOLLARS
 SINCE FILE
 TOTAL

 ENTRY
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 FULL ESTIMATED COST
 178.36
 178.36

FILE 'CAPLUS' ENTERED AT 14:47:07 ON 18 AUG 2008 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2008 AMERICAN CHEMICAL SOCIETY (ACS)

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FILE COVERS 1907 - 18 Aug 2008 VOL 149 ISS 8 FILE LAST UPDATED: 17 Aug 2008 (20080817/ED)

Caplus now includes complete International Patent Classification (IPC) reclassification data for the second quarter of 2008.

Effective October 17, 2005, revised CAS Information Use Policies apply. They are available for your review at:

http://www.cas.org/legal/infopolicy.html

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L4 ANSWER 1 OF 9 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2007:410774 CAPLUS DOCUMENT NUMBER: 146:421985

TITLE: Preparation of isotopically substituted (deuterated)

(fused) imidazopyridines for the treatment of

gastrointestinal disorders

INVENTOR(S): Kohl, Bernhard; Zimmermann, Peter Jan; Zech, Karl;
Buhr, Wilm; Palmer, Andreas; Brehm, Christof; Chiesa,
Maria Vittoria; Kromer, Wolfdang; Postius, Stefan;

Simon, Wolfgang-Alexander; Holst, Hans Christof

PATENT ASSIGNEE(S): Altana Pharma AG, Germany

SOURCE: PCT Int. Appl., 62pp.
CODEN: PIXXD2

DOCUMENT TYPE: Patent
LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

FAMILY ACC. NUM. COU PATENT INFORMATION:

GΙ

	PATENT NO.					KIN	D	DATE								D	ATE	
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	WO	2007																
		W:	ΑE,	AG,	AL,	AM,	ΑT,	AU,	ΑZ,	BA,	BB,	BG,	BR,	BW,	BY,	ΒZ,	CA,	CH,
			CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,
			GE,	GH,	GM,	HN,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KM,	KN,	KP,
			KR,	KZ,	LA,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	LY,	MA,	MD,	MG,	MK,	MN,
			MW,	MX,	MY,	MZ,	NA,	NG,	NI,	NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RS,
			RU,	SC,	SD,	SE,	SG,	SK,	SL,	SM,	SV,	SY,	TJ,	TM,	TN,	TR,	TT,	TZ,
			UA,	UG,	US,	UZ,	VC,	VN,	ZA,	ZM,	ZW							
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			IS,	IT,	LT,	LU,	LV,	MC,	NL,	PL,	PT,	RO,	SE,	SI,	SK,	TR,	BF,	BJ,
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			GM,	KE,	LS,	MW,	MZ,	NA,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,	AZ,	BY,
			KG.	KZ,	MD.	RU.	TJ.	TM										
	EP	1934	215			A1		2008	0625		EP 2	006-	7936	74		2	0060	920
		R:	AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,	FI,	FR.	GB,	GR,	HU,	IE,
								LV,										
				HR.														
PRIOR	ITY	APP									EP 2	005-	1087	64		A 2	0050	922
											EP 2							

W 20060920

WO 2006-EP66544
OTHER SOURCE(S): MARPAT 146:421985

$$R^3$$
 R^4
 R^5
 R^6
 R^2
 R^1
 R^9
 R^9

AB Title compds. [I; RI = H, alkyl, cycloalkyl, cycloalkylalkyl, alkoxy, alkoxyalkyl, alkoxycarbonyl, alkenyl, alkynyl, fluoroalkyl, hydroxyalkyl, R2 = H, alkyl, cycloalkyl, cycloalkylalkyl, alkoxycarbonyl, hydroxyalkyl, halo, alkenyl, alkynyl, fluoroalkyl, cycnomethyl, R3 = H, halo, alkyl, fluoroalkyl, cycleH, alkoxycarbonyl, hydroxyalkyl, alkoxyalkyl,

fluoroalkoxyalkyl, etc.; R4, R5 = H, R6 = (substituted) Ph; or R4R5 = CHR7CHR8; R7, R8 = H, OH, alkoxy, cycloalkylalkoxy, cycloalkylalkoxy, alkoxyalkoxy, fluoroalkoxy, hydroxyalkoxy, etc.; or R4 = H, R5R6 = Q1; Z = CHR11, CHR11CHR12; R9 = H, alkyl, hydroxyalkyl, alkoxy, alkenyloxy, aryloxy, etc.; R10 = H, alkyl, alkoxy, alkoxycarbonyl, halo, CF3, OH; R11, R12 = H, alkyl, alkenyl, OH, alkoxy, alkylcarbonylamino, etc.; X = O, NH; ≥ 1 of the H atoms of R1-R6 or of the core structure is replaced with D], were prepared Thus, Me 8 - [(2,6 - dimethylphenyl)] diducteromethylamino] -2,3 - dimethylphenyl) iducteromethylamino was heated 1 h with ethanolamine to give 73% 8 - [(2,6 - dimethylphenyl)] diducteromethylamino] -N (-2-hydroxyethyl) -2, -3

dimethylphenyldideuteromethylamino]-N-(2-hydroxyethyl)-2,3dimethylimidazo-6-carboxamide. The latter inhibited H+/K+-ATPase with -lg IC50 = 6.0.

IT 934248-01-8P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(claimed compound; preparation of isotopically substituted (deuterated) (fused) imidazopyridines for the treatment of gastrointestinal disorders)

RN 934248-01-8 CAPLUS

CN Imidazo[1,2-a]pyridine-6-carboxamide, 8-[[(2,6-dimethylphenyl)methyld2]amino]-N-(2-hydroxyethyl)-2,3-dimethyl- (CA INDEX NAME)

REFERENCE COUNT: 4 THERE ARE 4 CITED REFERENCES AVAILABLE FOR THIS
RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 2 OF 9 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2006:1173242 CAPLUS

DOCUMENT NUMBER: 145:489255

TITLE: Preparation of mutual prodrug compounds for use as

antiinflammatory agents with gastrointestinal
protective activity

INVENTOR(S): Brehm, Christof; Klein, Thomas; Buhr, Wilm; Chiesa,
Maria Vittoria; Palmer, Andreas; Zimmermann, Peter
Jan; Simon, Wolfdang-Alexander; Kromer, Wolfdangan

Postius, Stefan; Grundler, Gerhard

PATENT ASSIGNEE(S): Altana Pharma A.-G., Germany

SOURCE: PCT Int. Appl., 70pp.

CODEN: PIXXD2
DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

GI

	PATENT NO.					KIN	D	DATE			APPL		ION :			D	ATE	
	WO	2006	1173	 15		A1	_	2006	1109		WO 2					2	0060	426
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			GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KM,	KN,	KP,	KR,
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			VN,	YU,	ZA,	ZM,	zw											
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			IS,	IT,	LT,	LU,	LV,	MC,	NL,	PL,	PT,	RO,	SE,	SI,	SK,	TR,	BF,	ВJ,
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	ΕP	1879	891			A1		2008	0123		EP 2	006-	7548	65		2	0060	426
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			IS,	IT,	LI,	LT,	LU,	LV,	MC,	NL,	PL,	PT,	RO,	SE,	SI,	SK,	TR,	AL,
			BA,	HR,	MK,	YU												
PRIOR	IT	APP	LN.	INFO	. :						EP 2	005-	1035	81		A 2	0050	429
											WO 2	006-	EP61	850		W 2	0060	426
OTHER	SC	URCE	(S):			MAR	PAT	145:	4892	55								

AB The invention concerns A-Y-X-z-C(0)0-B (A is derived from ACO2H having antipyretic, analgesic, antiphlogistic and/or antiinflammatory properties; B is derived from HOB that are potassium competitive acid blockers; X = bond or linker (e.q. (un)substituted -(CH2)nOm(CH2)pOq(CH2)r (n = 1-7; m =

0, 1; p = 0.7; q = 0, 1; r = 0.7); Y = -C(0)0 with A attached to the carbonyl carbon; z = bond, -O-, -CHR1- or -NR1- (R1 = H or C1-4 alkyl); or X, Y and z together form a bond; addnl. details including provisos are given in the claims; e.g. (S)-2-(6-methoxynaphthalen-2-y1)propionic acid 3-[[[(7R.8R.9R)-2.3-dimethyl-7-(2-methoxyethoxy)-9-phenyl-7.8.9.10tetrahydroimidazo[1,2-h][1,7]naphthyridin-8-yl]oxy]carbonyl]propyl ester (shown as I)) and their salts. The compds. are prodrugs and exhibit in the human and/or animal body antipyretic, analgesic, antiphlogistic and/or antiinflammatory activity as well as gastric acid secretion inhibiting and therefore gastro and intestinal protective activity. Although the methods of preparation are not claimed, prepns. and/or characterization data for 23 examples of I and similar compds. are included. For example, I was prepared from (S)-2-(6-methoxynaphthalen-2-yl)propionic acid and 4-hydroxybutyric acid (7R, 8R, 9R) -2, 3-dimethyl-7-(2-methoxyethoxy) -9-phenyl-7, 8, 9, 10tetrahydroimidazo[1,2-h][1,7]naphthyridin-8-yl ester in THF using DMAP and toluenesulfonyl chloride. Data are provided for the inhibition of gastric acid secretion by 2 examples of I or similar compds. and for inhibition of COX-1/2 by 11 examples of I or similar compds.

IIT 248919-64-4, 2,3-Dimethyl-8-[(2,6-dimethylbenzyl)amino]-6-[N-(2-hydroxyethyl)aminoarbonyl]imidazo[1,2-a]pyridine
RL: RCT (Reactant); RRCT (Reactant or reagent)

(preparation of mutual prodrug compds. for use as antiinflammatory agents with gastrointestinal protective activity)

RN 248919-64-4 CAPLUS CN Imidazo[1,2-a]pvrid

Imidazo[1,2-a]pyridine-6-carboxamide, 8-[[(2,6dimethylphenyl)methyl]amino]-N-(2-hydroxyethyl)-2,3-dimethyl- (CA INDEX NAME)

$$\begin{array}{c} \text{Me} & \text{CH}_2\\ \text{CH}_2\\ \text{NH} \\ \text{HO-CH}_2\text{-CH}_2\text{-NH-C} \\ \text{O} \end{array}$$

REFERENCE COUNT:

2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT L4 ANSWER 3 OF 9 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2005:570894 CAPLUS

DOCUMENT NUMBER: 143:83527

TITLE: Crystalline forms of 2,3-dimethyl-8-(2,6-

dimethylbenzylamino)-N-hydroxyethylimidazo[1,2-

a]pyridine-6-carboxamide mesylate salt

INVENTOR(S): Lilljequist, Lars; Lindkvist, Maria; Nordberg, Peter;

Pettersson, Ursula; Sebhatu, Tesfai

PATENT ASSIGNEE(S): Astrazeneca AB, Swed. SOURCE: PCT Int. Appl., 66 pp.

CODEN: PIXXD2
DOCUMENT TYPE: Patent

LANGUAGE: English FAMILY ACC. NUM. COUNT: 1

FAMILY ACC. NUM. COU PATENT INFORMATION:

PA:	TENT	NO.			KIN	D	DATE				LICAT				D.	ATE	
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											, SC,						
		TJ.	TM.	TN.	TR.	TT,	TZ.	UA.	UG,	US	. UZ.	VC.	VN.	YU,	ZA.	ZM.	ZW
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	2004						2008	0417									
CA	2549	144			A1		2005	0630		CA	2004-	2549	144		2	0041	216
EP	1697	360			A1		2006	0906		EP	2004-	8090	82		2	0041	216
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		BA,	HR,	IS,	YU												
CN	1894	246			A		2007	0110			2004-					0041	216
BR	2004	0176	40		A		2007	0327			2004-					0041	
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IN	2006	DN03	006		A		2007	0803			2006-						
MX	2006	PA06	708		A		2006	0818		MX	2006-	PA67	80		2	0060	613
	2007						2007				2006-						
NO	2006	0033	09		A		2006	0914		NO	2006-	3309			2	0060	717
IORIT:	Y APP	LN.	INFO	. :						SE	2003-	3451			A 2	0031	218
										WO	2004-	SE19	09		W 2	0041	216

AB The present invention relates to novel crystalline forms of 2,3-dimethyl-8-(2,6-dimethylbenzylamino)-N-hydroxyethylimidazo[1,2-a]pyridine-6-carboxamide mesylate salt (I) and to mixture thereof. Further, the present invention also relates to processes for obtaining them, the use of the compds. for the treatment of gastrointestinal disorders, and pharmaceutical compns. containing them. 2,3-Dimethyl-8-(2,6-dimethylbenzylamino)-N-hydroxyethylimidazo[1,2-a]pyridine-6-carboxamide was treated with methanesulfonic acid in EtOH to give crystals of I Form A. The compound was characterized by x-ray crystallog.

IT 855998-67-3P

RL: PRP (Properties); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(crystalline forms of (dimethylbenzylamino)hydroxyethylimidazopyridinecarbox amide)

RN 855998-67-3 CAPLUS

CN Imidazo[1,2-a]pyridine-6-carboxamide, 8-[[(2,6-

```
dimethylphenyl)methyl]amino]-N-(2-hydroxyethyl)-2,3-dimethyl-,
methanesulfonate (1:1) (CA INDEX NAME)
```

CM 1

CRN 248919-64-4 CMF C21 H26 N4 O2

CM 2

CRN 75-75-2 CMF C H4 03 S

IT 248919-64-4

RL: RCT (Reactant); THU (Therapeutic use); BIOL (Biological study); RACT (Reactant or reagent); USES (Uses)

(crystalline forms of (dimethylbenzylamino)hydroxyethylimidazopyridinecarbox amide)

RN 248919-64-4 CAPLUS CN Imidazo[1.2-a]pyrid

Imidazo[1,2-a]pyridine-6-carboxamide, 8-[[(2,6dimethylphenyl)methyl]amino]-N-(2-hydroxyethyl)-2,3-dimethyl- (CA INDEX NAME)

REFERENCE COUNT:

THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 4 OF 9 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2005:409313 CAPLUS

DOCUMENT NUMBER: 142:457095

TITLE: Imidazo [1,2-a] pyridine derivatives for the treatment

of silent gastro-esophageal reflux

INVENTOR(S): Fernstroem, Paula; Hasselgren, Goeran

PATENT ASSIGNEE(S): Astrazeneca AB, Swed. SOURCE: PCT Int. Appl., 40 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PA	TENT				KIN	D	DATE			APPL					D.	ATE	
WO	2005				A1		2005	0512							2	0041	103
	W:	AE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,
		CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,
		GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC,
		LK,	LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NA,	NI,
		NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SY,
		TJ,	TM,	TN,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VC,	VN,	YU,	ZA,	ZM,	ZW
	RW:	BW,	GH,	GM,	KE,	LS,	MW,	MZ,	NA,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,
		AZ,	BY,	KG,	KZ,	MD,	RU,	TJ,	TM,	AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,
		EE,	ES,	FI,	FR,	GB,	GR,	HU,	IE,	IS,	IT,	LU,	MC,	NL,	PL,	PT,	RO,
		SE,	SI,	SK,	TR,	BF,	ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,
		NE,	SN,	TD,	TG												
AU	2004	2853	94		A1		2005	0512		AU 2	004-	2853	94		2	0041	103
CA	2544	325			A1		2005	0512		CA 2	004-	2544	325		2	0041	103
EP	1682	133			A1		2006	0726		EP 2	004-	8002	52		2	0041	103
	R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PT,
		IE,	SI,	LT,	LV,	FI,	RO,	CY,	TR,	BG,	CZ,	EE,	HU,	PL,	SK,	HR,	IS
CN	1874	772			A		2006	1206		CN 2	004-	8003	2415		2	0041	103
IN	2006	DN01	943		A		2007	0803		IN 2	006-	DN19	43		2	0060	410
NO	2006	0025	70		Α		2006	0803		NO 2	006-	2570			2	0060	602
PRIORIT	Y APP	LN.	INFO	. :						US 2	003-	5171	25P		P 2	0031	103
										WO 2	004-	SE15	89		W 2	0041	103
OTHER S	OURCE	(S):			MAR	PAT	142:	4570	95								

AB The present invention relates to a new method of treatment of sleep disturbance due to silent gastro-esophageal reflux. The invention further

Ι

relates to the use of potassium-competitive acid blockers (P-CAB's) which inhibit the enzyme responsible for gastric acid secretion (H+KH-ATBase). In particular, the present invention relates to the use of certain imidazo (1,2-a)pyridines derivs. (I wherein R1 = H, Me or Bt: R2 = Me or Et; R3 and R4 = H, C1-6 alkyl, hydroxylated C1-6 alkyl or halogen; R5 = H or halogen; R6 and R7 = H, C1-6 alkyl, hydroxylated C1-6 alkyl or C1-6 alkyl or C1-6 alkyl and X = NH or O) in said treatment.

IT 248919-64-4
RL: PAC (Pharmacological activity); THU (Therapeutic use); BIOL

(Biological study); USES (Uses) (imidazo(a)pyridine derivs. for treatment of silent gastro-esophageal reflux and sleep disturbances in relation to potassium-competitive acid secretion blockade)

RN 248919-64-4 CAPLUS

CN Imidazo[1,2-a]pyridine-6-carboxamide, 8-[[(2,6-dimethylphenyl)methyl]amino]-N-(2-hydroxyethyl)-2,3-dimethyl- (CA INDEX NAME)

$$\begin{array}{c} \text{Me} & \text{CH}_2 \\ \text{CH}_2 \\ \text{NH} \\ \text{HO-CH}_2\text{-CH}_2\text{-NH-C} \\ \text{O} \end{array}$$

REFERENCE COUNT:

THERE ARE 8 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 5 OF 9 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2004:1059201 CAPLUS 142:32977

DOCUMENT NUMBER:

TITLE: Pharmaceutical combinations of a proton pump inhibitor and a compound which modifies gastrointestinal

motility

Zimmermann, Peter Jan; Chiesa, M. Vittoria; Palmer, INVENTOR(S): Andreas; Brehm, Christof; Klein, Thomas;

> Senn-Bilfinger, Joerg; Simon, Wolfgang-Alexander; Kromer, Wolfgang; Grundler, Gerhard; Hanauer, Guido;

Buhr, Wilm; Postius, Stefan Altana Pharma A.-G., Germany

PATENT ASSIGNEE(S): SOURCE: PCT Int. Appl., 102 pp.

CODEN: PIXXD2 DOCUMENT TYPE: Patent

LANGUAGE: English FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

		TENT I						DATE					ION				ATE		
		2004															0040	526	
		W:	ΑE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BW,	BY,	ΒZ,	CA,	CH,	
			CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FΙ,	GB,	GD,	
			GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC,	
			LK,	LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NA,	NI,	
			NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SY,	
			TJ,	TM,	TN,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VC,	VN,	YU,	ZA,	ZM,	ZW	
		RW:	BW,	GH,	GM,	KE,	LS,	MW,	MZ,	NA,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,	
			AZ,	BY,	KG,	KZ,	MD,	RU,	TJ,	TM,	AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	
			EE,	ES,	FI,	FR,	GB,	GR,	HU,	ΙE,	IT,	LU,	MC,	NL,	PL,	PT,	RO,	SE,	
			SI,	SK,	TR,	BF,	ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,	NE,	
				TD,															
		2004																	
		2526																	
	ΕP	1644	043			A1		2006	0412		EP 2	004-	7416	58		2	0040	526	
		R:	AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PT,	
								RO,											HR
	JΡ	2006	5282	31		T		2006	1214		JP 2	006-	5302	22		2	0040	526	
		2005																	
		2006																	
	NO	2005	0059	68		A		2005	1215		NO 2	005-	5968			2	0051	215	
PRIOR	RIT	APP:	LN.	INFO	. :						EP 2	003-	1187	5		A 2	0030	527	
											EP 2	004-	1023	04		A 2	0040	525	
											WO 2	004-	EP50	936		W 2	0040	526	
AB	The	e inv	enti	on r	elat	es t	o th	e co	mbin	atio	n of	cer	tain	act.	ive	comp	ds.	from	

Α the acid pump antagonist class and compds. which modify gastrointestinal motility. The acid pump antagonist class is selected from a tricyclic imidazopyridine and the gastrointestinal motility modifier is selected from a 5-HT-(partial)-agonist/antagonist.

248919-64-4

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (pharmaceutical combinations of proton pump inhibitor and modifier of gastrointestinal motility)

248919-64-4 CAPLUS

Imidazo[1,2-a]pyridine-6-carboxamide, 8-[[(2,6dimethylphenyl)methyl]amino]-N-(2-hydroxyethyl)-2,3-dimethyl- (CA INDEX NAME)

16

REFERENCE COUNT:

THERE ARE 16 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 6 OF 9 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2003:913040 CAPLUS

DOCUMENT NUMBER: 139:375018

TITLE: Combinations containing proton pump inhibitors for the

treatment of airway disorders

INVENTOR(S): Hanauer, Guido; Kromer, Wolfgang; Postius, Stefan;

Simon, Wolfgang-Alexander PATENT ASSIGNEE(S): Altana Pharma A.-G., Germany

SOURCE: PCT Int. Appl., 21 pp.

CODEN: PIXXD2 DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

	TENT N						DATE				LICAT					ATE	
WO	20030	9496	57		A2		2003	1120									
		IS,	JP,		LT,						, DZ, , NZ,						
	RW:	AM, DK,	AZ, EE,	BY,	KG,						, AT,						
AU	20032				A1		2003	1111		AH	2003-	2277	1.0		2	0030	503
	24842																
	15060																
											, IT,						
		IE,	SI,	LT,	LV,	FI,	RO,	MK,	CY,	AL	, TR,	BG,	CZ,	EE,	HU,	SK	
BR	20030	0980	8(A		2005	0301		BR	2003-	9808			2	0030	503
CN	16528 20055	22			Α		2005	0810		CN	2003-	8104	00		2	0030	503
JP	20055	2841	18		T		2005	0922		JΡ	2004-	5030	50		2	0030	503
	2004M										2004-						
ZA	20040	0789	96		Α												
	2004P										2004-						
	20050																
	20040				A		2004	1206			2004-						
PRIORIT:	Y APPL	N. 1	NFO	. :							2002-						
										WO	2003-	EP46	53	1	W 2	0030	503

A method for treating airway disorders comprises a reversible proton pump inhibitor and an airway therapeutic to be taken simultaneously (as a fixed oral combination) or in succession (one directly after the other or else within a relatively large time span). The reversible proton pump inhibitor is, e.g., Soraprazan or its salt, and the airway therapeutic is, e.g., Ciclesonide.

248919-64-4

RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses) (oral combination of reversible proton pump inhibitors and airway therapeutics for treatment of airway disorders)

248919-64-4 CAPLUS

CN Imidazo[1,2-a]pyridine-6-carboxamide, 8-[[(2,6dimethylphenyl)methyllaminol-N-(2-hydroxyethyl)-2,3-dimethyl- (CA INDEX

NAME)

$$\begin{array}{c} \text{Me} \\ \text{CH2} \\ \text{NH} \\ \text{HO-CH2-CH2-NH-C} \\ \text{O} \\ \end{array}$$

L4 ANSWER 7 OF 9 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2002:637503 CAPLUS

DOCUMENT NUMBER: 137:190728

TITLE: Novel modified release formulation containing

carboxamide derivatives for inhibition of secretion of gastric acid

INVENTOR(S): Juppo, Anne

PATENT ASSIGNEE(S): Astrazeneca Ab, Swed.
SOURCE: PCT Int. Appl., 35 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent
LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2 PATENT INFORMATION:

												LICAT						
												2002-						
		W:	ΑE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB	, BG,	BR,	BY,	BZ,	CA,	CH,	CN,
			CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC	, EE,	ES,	FI,	GB,	GD,	GE,	GH,
			GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE	, KG,	KP,	KR,	ΚZ,	LC,	LK,	LR,
			LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN	, MW,	MX,	MZ,	NO,	NZ,	OM,	PH,
			PL,	PT,	RO,	RU,	SD,	SE,	SG,	SI,	SK	, SL,	ТJ,	TM,	TN,	TR,	TT,	TZ,
			UA,	UG,	US,	UZ,	VN,	YU,	ZA,	ZM,	ZW							
		RW:										, TZ,						
												, IT,						
												, GW,						
C.	A 2	4345	542			A1		2002	0822		CA	2002-	2434.	542		2	0020	208
												2002-						
E												2002-						
		R:										, IT,			NL,	SE,	MC,	PT,
			IE,	SI,	LT,	LV,	FΙ,	RO,	MK,	CY,	AL	, TR						
C	N 1	.491	105			A		2004	0421		CN	2002-	8049	06		2	0020	208
C	N 1	491:	104			A		2004	0421		CN	2002- 2002-	8049	14		2	0020	208
J	P 2	2004	51870	08		т		2004	0624		JP	2002-	5639	14		2	0020	208
N	Z 5	2699	93			A		2005	0128		NZ	2002-	5269	93		2	0020	208
A	T 3	248	/1			T						2002-						
P	TI	3681	006			T						2002-						
		261				T3 A						2002-						
												2003-						
						A1						2003-						
PRIORI						AI		2008	0522			2007-						
FKIOKI	11	neri	DIN	TIME	• •							2001-						
												2001-					0010	
												2002-						

OTHER SOURCE(S): MARPAT 137:190728 GI

IΤ

Т

- AB A multiparticulate (particle size < 300 µm), modified-release solid dispersion formulation comprises (i) a drug substance having a pH-dependent solubility, i.e., compound I (RI = H, Me, Et; R2 = Me, Et; R3, R4
 - H, C1-6 alkyl, hydroxylated C1-6 alkyl, halogen; R5 = H, halogen; R6, R7 = H, C1-6 alkyl, hydroxylated C1-6 alkyl, C1-6 alkoxy-substituted C1-6 alkyl; X = NH, O) or a pharmaceutically acceptable salt thereof; (ii) a hydrophobic matrix former which is a water-insol., non-swelling amphiphilic lipid; and (iii) a hydrophilic matrix former which is a meltable, water-soluble excipient. The weight ratio of hydrophobic matrix former/hydrophilic matrix former is ≥1 and the particle size is less than 300 µm. Also a unit dosage form of the compound I, as well as a process for its preparation, and the use of the formulation and unit dosage form for inhibiting the secretion of gastric acid are described. For example, multiparticulate, modified-release formulation was prepared by dissolving 1 g of 2,3-dimethyl-8-(2-ethyl-6-methylbenzylamino)imidazo[1,2a]pyridine-6-carboxamide mesylate in a melt of 4 g myristic acid at 90° and adding 2 g of polyethylene glycol 4000 (PEG 4000) into the melt. The melted mixture was atomized at 90° and the particles were collected into a vessel which was kept on ice. The resulted particles were spherical and < 300 μm in size. The amount of 3 q of particles were blended with 5.85 g microcryst, cellulose and 0.016 g sodium stearyl fumarate and compressed into 450 mg tablets. The dissoln, of tablets was 52-56% in 3 h. 248919-64-4
- RL: THU (Therapeutic use); BIOL (Biological study); USES (Uses)
 (controlled-release formulation containing imidazopyridine carboxamide
 derivs. for inhibition of gastric acid secretion)
 RN 248919-64-4 CAPLUS

 CON Inidayoli 2-alpuvidine-6-carboxamide 8-11(2.6-
 - $\label{limit_eq} $$ \operatorname{Imidazo}(1,2-a) \operatorname{pyridine-6-carboxamide}, \ \$-[[(2,6-dimethylphenyl)methyl] = -(2-hydroxyethyl)-2,3-dimethyl-NAME) $$ NAME) $$$

REFERENCE COUNT:

THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 8 OF 9 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 2002:185119 CAPLUS DOCUMENT NUMBER: 136:249369

TITLE: Process for preparing a substituted imidazopyridine

compound INVENTOR(S): Elman, Bjoern; Erback, Silke; Thiemermann, Eric

PATENT ASSIGNEE (S): SOURCE:

Astrazeneca AB, Swed. PCT Int. Appl., 27 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA:	TENT :	NO.			KIN)	DATE			APP	LICAT	ION	NO.			DATE	
WO	2002	0205	23		A1		2002	0314		wo	2001-	SE18	97			20010	905
	W:										, BG,						
											EE,						
											, KG,						
											MW.						
											, TJ.						
							ZW	,	~,	~~	,,	,	,	,		,,	,
	RW:							SD.	SI	S7	, TZ,	UG.	7W.	AT.	BE	. CH.	CY.
											, LU,						
											, ML,						,
CA	2419	764	02 /	00,	A1	011,	2002	0314	027	CA.	2001-	2419	764	0117		20010	905
AII	2001	08459	9.4		A		2002	0322		AII	2001- 2001-	8459	4			20010	905
EP	1317	455			A1		2003	0611		EP	2001-	9636	65			20010	905
EP	1317	455			B1		2004	0804			2001-	,,,,,					
	R:	AT.	BE.	CH.	DE.	DK.	ES.	FR.	GB.	GR	, IT,	LI.	LU.	NL.	SE	MC.	PT.
		IE.	SI.	LT.	LV.	FI.	RO.	MK.	CY.	AL	TR		,		-	,,	
BR	2001	01360	02	,	A	,	2003	0715	,	BR	2001-	1360	2			20010	905
HU	2003	0022	77		A2		2003	1028		HU	2003-	2277				20010	905
HU	2003	0022	77		A3		2003	1229									
HU	2254	59			В1		2006	1228									
JP	2004	5083	71		T		2004	0318		JP	2002-	5251	44			20010	905
AT	2726	37			T		2004	0815		ΑT	2001-	9636	65			20010	905
NZ	5243	02			A		2004	0827		NZ	2001-	5243	02			20010	905
PT	1317	455			T		2004	1130		PΤ	2001-	9636	65			20010	905
EE	2003	0009	0		A		2004	1215		EE	2003-	90				20010	905
ES	2223	906			Т3		2005	0301		ES	2001-	9636	65			20010	905
CZ	2949	57			В6		2005	0413		CZ	2003-	643				20010	905
AU	2001	28459	94		B2		2005	1215		AU	2001-	2845	94			20010	905
RU	2275	372			C2		2006	0427		RU	2003-	1049	87			20010	905
ZA	2003	0011	71		A		2004	0318		ZA	2003-	1171				20030	212
IN	2003	MN00:	220		A		2006	0505		IN	2003-	MN22	0			20030	214
MX	2003	PA019	941		A		2003	0624		MX	2003-	PA19	41			20030	305
NO	2003	0010	46		A		2003	0505		NO	2003-	1046				20030	306
NO	3242	52			B1		2007	0917									
KR	7704	78			B1		2007	1026		KR	2003-	7033	11			20030	306
US	2004	00390	013		A1		2004	0226		US	2003-	3638	06			20030	627
US	6900	324			B2		2005	0531									
HK	1054	388			A1		2005	0408		HK	2003-	1066	57			20030	916
US	2006	0063	797		A1		2006	0323		US	2005-	1073	52			20050	414
RIORIT:	Y APP	LN. :	INFO	.:						SE	2000-	3186			A	20000	907
										WO	2001-	SE18	97		W	20010	905
										US	2003-	3638	06		A1	20030	627
THER SO	DURCE	(S):			MAR	PAT	136:	2493	69		, IT, , TR, , TR, , TR 2002-2001-2003- 2001-2001-2001-2003- 2003-2003-2003-2003-2003-2003-2003-2003						

- AB Present invention provides a new process for large-scale preparation of substituted imidazopyridine compound of formula (1), wherein R1 = C1-6 alkoxy or NH2 group, comprising the step of reacting a compound of formula (11) with a 3-halo-2-butanone compound in cyclohexanone. Thus, 5.1 g 5,6-diaminonicotinic acid Me ester, 50 mL cyclohexanone, and 3.9 mL bromobutanone were agitated at 100° for 2.5 h to give Me 8-amino-2,3-dimethylimidaco(1,2-a]pyridine-6-carboxylate.
- IT 248919-64-4P RL: IMF (Industrial manufacture); PREP (Preparation)
- (process for preparing a substituted imidazopyridine compound)
 RN 248919-64-4 CAPLUS
- CN Imidazo[1,2-a]pyridine-6-carboxamide, 8-[[(2,6-
- dimethylphenyl)methyl]amino]-N-(2-hydroxyethyl)-2,3-dimethyl- (CA INDEX NAME)

REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L4 ANSWER 9 OF 9 CAPLUS COPYRIGHT 2008 ACS on STN

ACCESSION NUMBER: 1999:708770 CAPLUS

DOCUMENT NUMBER: 131:322617

TITLE: Preparation of imidazopyridines which inhibit gastric

acid secretion

INVENTOR(S): Amin, Kosrat; Dahlstrom, Michael; Nordberg, Peter;

Starke, Ingemar PATENT ASSIGNEE(S): Astra AB, Swed.

SOURCE: PCT Int. Appl., 77 pp.

CODEN: PIXXD2
DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 2

PATENT INFORMATION:

	ENT				KIN	D	DATE			APP	LICAT	ION	NO.		D.	ATE	
	9955										 1999-					9990	
	W:	AE,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG	, BR,	BY,	CA,	CH,	CN,	CU,	CZ,
											, GM,						
		JP,	KE,	KG,	KP,	KR.	KZ,	LC,	LK,	LR	, LS,	LT,	LU,	LV,	MD,	MG,	MK,
		MN.	MW.	MX.	NO.	NZ.	PL.	PT.	RO.	RU	, SD,	SE.	SG.	SI.	SK.	SL,	TJ.
											, ZA,						
	RW:										, ZW,		BE,	CH,	CY,	DE,	DK,
											, NL,						
TW	4904	66			В		2002	0611		TW	, TD, 1999- 1999- 1999- 1999- 1999-	8810	6129		1	9990	416
TW	2501	59			В		2006	0301		TW	1999-	8810	6128		1	9990	416
CA	2329	922			A1		1999	1104		CA	1999-	2329	922		1	9990	423
CA	2329	922			C		2006	0411									
AII	9943	007			A		1999	1116		AII	1999-	4300	7		1	9990	423
ΑU	7691	90			B2		2004	0122									
BR	9909	996			Α		2000	1226		BR	1999-	9996			1	9991	423
EP	1073	657			A1		2001	0207		EP	1999-	9470	3.8		1	9990	423
EP	1073	657			B1		2005	1207							-	,,,,	
	B.	AT.	BE.	CH.	DE.	DK.	ES.	FR.	GB.	GR	, IT,	LT.	LII.	NT.	SE.	MC.	PT.
		TE,	SI	I.T	LV	FT.	RO,	CY	02,	010	,,	,	20,	,	02,	110,	/
TD	2000	0314	۵.,	,	Т2	,	2001	0321		TD	2000- 2000- 2001- 2000- 2001- 2001- 2001- 2000- 2000- 2000- 2000- 2000-	3149			1	aaan	423
TR	2000	0317	6		T2		2001	0321		TR	2000-	3176			1	9990	423
HII	2001	0017	25		Δ2		2001	1128		HII	2001-	2425			1	9990	423
HII	2001	0024	25		A3		2002	1228			2001	_ 1_0			-	,,,,	
FF	2001	0024	4		71.0		2002	0/15		FF	2000-	661			1	aaan	123
FF	4916	0000	4		R1		2002	1015			2000	004			_	,,,,	425
TD	2002	5130	25		T		2007	0500		TD	2000-	5/150	65		1	aaan	123
TD	2602	034	23		D2		2002	0000		OF	2000-	3430	0.5			2220	125
TD	2001	0261	2		T2		2003	0601		TD	2001	2612			1	0000	122
TD	2001	0201	6		T2		2002	0621		TD	2001-	2720			1	0000	122
TIV.	2001	67	0		DC.		2002	1015		CZ	2001-	2002			1	0000	423
NZ	2923	20			3		2003	1013		UZ NE	1000	5902	20		1	9990	423
NZ	2076	33			D.C		2004	0130		NZ OF	1999-	2076	39		1	9990	423
CZ	2939	071			86		2004	0915		CZ	2000-	3981	10		1	9990	423
RU	2238	2/1			7.0		2004	1020		KU	2000-	12/0	19		1	9990	423
EP	1491	542			AZ		2004	1229		EP	2004-	2309	U		1	9990	423
EP	1491	542			A.3		2005	0105									
EP	1491	542		011	BI	DI	2005 2007 ES.	0905	on.	on	T. III			***	0.0		D.M.
	K:																
		IE,	SI,	LT,	LV,	E.T.	RO,	MK,	CY								
	1491	543			A1		2004	1229		EP	2004-	2309	T		1	9990	423
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	R:									GR	, IT,	LΙ,	LU,	ΝL,	SE,	MC,	PT,
		IE,	SI,	LT,	LV,	FΙ,	RO,	MK,	CY								
AΤ					Т		2005	1215					4 R		1	999N	423
	3121	01								711	1999- 1999-	3470			_		

ES 2252975	Т3	20060516	ES	1999-947038		19990423
SK 285768	В6	20070706	SK	2000-1492		19990423
PL 195000	B1	20070731	PL	1999-343801		19990423
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AT 372340	T	20070915	AT	2004-23091		19990423
US 6313137	B1	20011106	US	1999-319973		19990614
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MX 2000PA10239	A	20010405		2000-PA10239		20001019
NO 2000005450	A	20001222	NO	2000-5450		20001027
NO 317262	B1	20040927				
HK 1071140	A1	20080215	HK	2005-103979		20010612
HK 1033317	A1	20060630		2001-104064		20010613
HK 1036984	A1	20050429		2001-107857		20011108
PRIORITY APPLN. INFO.:			SE	1998-1526	A	19980429
				1999-947037		19990423
				1999-947038	A3	19990423
				1999-SE663	W	19990423
			HK	2001-104026	A3	20010612
OTHER SOURCE(S):	MARPAT	131:322617				

R6 N N N R2 X R3 R5

GI

AB The title compds. [I; R1 = H, Me, CH2OH; R2 = Me, Et; R3 = H, alkyl, halo, etc.; R4 = H, alkyl, halo, etc.; R5 = H, halo; R6, R7 = H, alkyl, hydroxylated alkyl, etc.; X = NH, O] which inhibit exogenously or endogenously stimulated gastric acid secretion (no data) and thus can be used in the prevention and treatment of gastrointestinal inflammatory diseases, and for treatment or prophylaxis of conditions involving infection by Helicobacter pylori of human gastric mucosa, were prepared Thus, reacting Et 2,3-dimethyl-8-(2-ethyl-6-methylbenzylamino)-imidazo[1,2-a]pyridine-6-carboxylate with propylamine in the presence of a cat. amount of NaCN in MeOH afforded 42% I [R1 = R2 = R4 = Me; R3 = Et; R5 = R7 = H; R6 = Pr]. In general, compds. I are effective at 5-1000 mg/day.

RL: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SFN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of imidazopyridines which inhibit gastric acid secretion) ${\tt RN} - 248919-64-4 - {\tt CAPLUS}$

CN Imidazo[1,2-a]pyridine-6-carboxamide, 8-[[(2,6-

т

REFERENCE COUNT: 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

(FILE 'HOME' ENTERED AT 14:46:31 ON 18 AUG 2008)

FILE 'REGISTRY' ENTERED AT 14:46:40 ON 18 AUG 2008

L1 STRUCTURE UPLOADED

L2 0 S L1 L3 3 S L1 FULL

CA SUBSCRIBER PRICE

FILE 'CAPLUS' ENTERED AT 14:47:07 ON 18 AUG 2008 L4 9 S L3 FULL

=> log y

COST IN U.S. DOLLARS
SINCE FILE TOTAL
ENTRY SESSION
FULL ESTIMATED COST 49.53 228.10

DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS) SINCE FILE TOTAL
ENTRY SESSION

-7.20

-7.20

STN INTERNATIONAL LOGOFF AT 14:47:44 ON 18 AUG 2008